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Ethanol formation from CO₂ hydrogenation at atmospheric pressure using Cu catalysts: Water as a key component

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ABSTRACT

The catalytic conversion of CO_2 can reduce its impact in the atmosphere and produce chemicals of industrial interest. Based on this, herein, ethanol formation from CO_2 hydrogenation with water was studied. Inspired by CO_2 electroreduction results, we show that ethanol can be formed at atmospheric pressure, using metallic Cu catalysts in the CO_2 hydrogenation with water steam ($CO_2 + H_2 + H_2O$) with selectivity of 84% and productivity of $\sim 2 \, \mu mol.gcat^{-1}.h^{-1}$ at 190 °C. When only H_2O is used (without H_2), the same trend was observed. To the best of our knowledge, ethanol is reported for the first time to be synthesized at atmospheric pressure, using only CO_2 and water as reactants in a thermocatalytic process. 1H NMR results showed that water (and deuterated water) hydrogenate CO_2 to form ethanol. CO-DRIFTS analyses revealed that water enhances the carbon-metal strengthening and this can explain why ethanol production is favored during the CO_2 hydrogenation.

1. Introduction

The atmospheric CO_2 concentration has increased by over 100 ppm in the last century, whereby more than 20 ppm has increased over the last ten years. This increase is mainly caused by the agriculture and industrial expansions that are essential for human needs, and therefore these activities cannot be quickly interrupted. The best strategy to reduce the impact of human activities is to convert CO_2 into desired products that can be reused by society. To this end, catalytic hydrogenation of CO_2 is one of the most promising applications due to its possibility of massively converting CO_2 into fuels and building blocks for the chemical industry. For example, the synthesis of chemicals with two or more carbons (C_{2+} compounds), as ethanol, is a desired reaction because it can both reduce the impact of this gas in the atmosphere and produce a chemical of high interest due to its broad applicability in the industry and our daily life.

Effective Cu-based catalysts have been used for CO_2 hydrogenation, such as modified industrial $Cu/ZnO/Al_2O_3$ [1,2], Cu/ZnO/M (M=Zr [3–5], Ce [6,7], Ti [8–10], Ga [11–13], Nb [14], and others [15,16]), and Pd-Cu [17,18], but mainly C_1 compounds (CO, CH_4 , and methanol) are reported as products. Such an approach has already been applied for

methanol synthesis on a large scale. George Olah Renewable Methanol plant in Reykjavik, Iceland, was the first company to produce ten tonmethanol/day from CO2 and H2 using heterogeneous catalysis and geothermal energy [19]. In contrast, the synthesis of higher alcohols, e. g., ethanol, from CO₂ hydrogenation is still far from a mature technology for industrial application because, differently from methanol synthesis, C-C coupling is needed to produce these compounds. The C-C coupling hampers the reaction pathway significantly, requiring catalysts that allow CO₂ and CO to be adsorbed strongly enough to prevent them from being desorbed as a product before being reacted, and not so strongly that the catalyst is not poisoned by the adsorption of carbon. Recently, many efforts have been dedicated to obtaining better catalytic performance in ethanol synthesis from CO₂ hydrogenation [20–26]. Applying noble metals like Pd and Iridium, high selective hydrogenation of CO2 to ethanol is achieved (> 90%), although CO₂ conversion is relatively low (< 10 %) [21,23,27]. Cu-, Co-, and Fe-based catalysts, otherwise, shows higher CO₂ conversion (> 30%) but the ethanol selectivity is compromised (< 40 %) [28-30]. Unique performance was found on Cu@Na-Beta, where ethanol was 100% selective at 300 $^{\circ}\text{C}$ and 1.3 MPa with 8% CO₂ conversion [25].

Although a lot of advances have been achieved on the ethanol

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synthesis from CO_2 hydrogenation, low productivity, low conversion, severe reaction conditions, and/or high costs involved in the process inhibit practical applications. One of these problems is using H_2 as a hydrogen source for CO_2 hydrogenation. CH_4 steam reforming is the primary source to produce H_2 , and the costs already involved in the production of H_2 make the conventional CO_2 hydrogenation to ethanol $(2CO_2+6\ H_2\leftrightarrow C_2H_5OH+3\ H_2O)$ economically unfeasible. Furthermore, the CH_4 is mainly obtained from fossil fuels, which impairs the sustainability of the process. A promising strategy is using H_2O to partially (or fully) suppress the use of H_2 as a hydrogen source [31]. CO_2 electroreduction already uses this approach, and it is well known that ethanol and other C_{2+} compounds can be synthesized from CO_2 and H_2O , applying metallic CU as the catalyst [32,33]. However, the high costs due to the overpotentials required for producing these chemicals also inhibit industrial applications.

In CO₂ hydrogenation, water is commonly avoided because it can sinter and oxidize the active phase [34,35], and only a few works study the impact of water on feed gas. He et al. [36] showed that higher alcohols (C₂-C₄) were synthesized at 80 bar (H₂/CO₂ = 3/1) and 140 $^{\circ}$ C applying Pt/Co₃O₄ in a batch reactor containing water as the solvent. They showed that water was responsible for promoting the reaction kinetically. Zhao et al. [37] performed CO₂ hydrogenation with water steam over a commercial Cu/ZnO/Al₂O₃ catalyst with the assistance of negative corona discharge plasma and showed that only ethanol was detected as the liquid product at 1 bar and 160 °C. Curiously, in the CO₂ hydrogenation with H₂ applying the industrial Cu/ZnO/Al₂O₃ catalyst, merely methanol and CO are produced (only C₁ compounds) [38-40]. Koishybay and Shantz [41] have shown that the water provides the hydroxyl group in the synthesis of methanol from partial oxidation of methane, which suggests that water plays an essential role in synthesizing alcohols. In the same direction, Wang et al. [42] reported strong evidence of methanol synthesis enhancement from CO2 hydrogenation over Cu-ZnO-ZrO₂ (CZZ) catalysts by increasing the methanol selectivity and the methanol yield, adding a certain amount of water to the feed gas. So, as commonly seen in the CO2 electroreduction approach, water in the reactional medium may play an essential role in synthesizing alcohols and C2 compounds via thermocatalysis although its impact is poorly investigated in literature and more evidence of the water impact in the products distribution of the CO₂ hydrogenation are still necessary.

Here, we investigated the impact of water on ethanol production from ${\rm CO_2}$ hydrogenation with water steam at atmospheric pressure, temperatures between 170 and 240 °C, and applying metallic Cu catalysts. Our results showed that the water steam can favor the formation of ethanol from ${\rm CO_2}$ while in its absence only CO and methanol were observed (${\rm C_1}$ compounds) on Cu catalysts. Thermodynamics were also discussed here. CO-DRIFTS indicate that water steam enhances the carbon-metal strengthening and this can explain why ethanol production is favored in the ${\rm CO_2}$ hydrogenation with water.

2. Experimental

2.1. Catalyst synthesis

2.1.1. Preparation of Cuprec

CuO powder was synthesized via the precipitation method. In a typical procedure, 3.80 g of $Cu(NO_3)_20.3$ H_2O (99 %, Sigma-Aldrich) were dissolved in 20 mL of deionized water. Then this solution containing Cu ions was slowly dripped into 400 mL of 0.1 M NaHCO₃ under constant stirring and at a temperature of 70 °C. The pH of the solution was controlled to be between 7 and 8. Decreases in pH due to the addition of the Cu precursor were compensated by the dropwise of a 0.5 M NaHCO₃. After completing the Cu precursor addition, the solution was aged for 2 h. Finally, the solid was separated from the supernatant by filtration, followed by washing (with at least 2 L of deionized water), dried at 80 °C for 12 h, and calcined for 5 h at 350 °C using a heating rate of 2 °C/min. Before the catalyst was used in the CO₂ hydrogenation, the

CuO powder was reduced in situ with H_2 , resulting in the metallic Cu powder, called $Cu_{\rm prec.}$.

2.1.2. Preparation of Cucalc

CuO powder was synthesized by direct calcination of Cu(NO₃)₂0.3 H₂O. Typically, 3.80 g of Cu(NO₃)₂0.3 H₂O (99 %, Sigma-Aldrich) were placed in a porcelain crucible, and the nitrate was calcined for 5 h at 350 $^{\circ}$ C using a heating rate of 2 $^{\circ}$ C/min. Before the catalyst was used in the CO₂ hydrogenation, the CuO powder was reduced in situ with H₂, resulting in the metallic Cu powder, named Cu_{calc}.

2.1.3. Preparation of Cucubes

 $\rm Cu_xO$ was synthesized according to the procedure proposed by Chang et al. [43]. In a typical procedure, 1 mL of 1.2 M CuSO₄ (CuSO₄, 99 %, Synth) was quickly added to a round bottom flask with 0.2 M $\rm C_6H_5Na_3O_7$ ($\rm C_6H_5Na_3O_70.2~H_2O$, 99%, Synth) at 25 °C and under vigorous stirring. After 5 min, 1 mL of 4.8 M NaOH (NaOH, 99 %, Aldrich) was added to the solution. The solution immediately turned turbid blue, indicating Cu(OH)₂ precipitation. After another 5 min, 1 mL of 1.2 M ascorbic acid was added to the solution as a reducing agent. The color of the solution rapidly turned from turbid blue to yellowish-brown, indicating the formation of Cu₂O. The solution was kept in a water bath and under vigorous stirring for another 30 min. The solid formed was filtered and washed with deionized water. Finally, the filtered solid was dried for 2 h at 80 °C. Before the catalyst was used in the CO₂ hydrogenation, the Cu_xO powder was reduced in situ with H₂, resulting in the metallic Cu powder, named Cu_{cubes}.

2.2. Catalyst characterization

Scanning electron microscopy (SEM) images were obtained in an Apreo SEM (ThermoFisher Scientific) with an acceleration voltage of 15 kV and an electron beam current of $0.4\ nA$.

The specific surface areas were measured in a Micromeritics ASAP 2020 instrument according to the traditional B.E.T method using N_2 physisorption at - 196 °C. The samples were pre-treated under vacuum for 2 h at 200 °C. Before the specific surface area be measured, the Cu_xO was reduced for 1 h at 300 °C.

The temperature-programmed reduction (TPR) and temperature-programmed desorption (TPD) analysis were performed on a Micromeritics ChemiSorb 2750 equipped with a thermal conductivity detector (TCD). First, 150 mg of catalyst were inserted in a U-tube reactor and pre-treated for 60 min at 300 °C in a flow of H₂ (40 mL/min). For the TPR experiment, the surface of the metallic Cu catalyst was oxidized to Cu $^+$ by flowing 10 % N₂O/Ar (25 mL/min) for 30 min at 30 °C followed by surface purging with Ar for additional 30 min at the same temperature. Afterward, the catalyst's surface was reduced during heating from 30° to 450°C in a 10 % H₂/Ar mixture at a flow rate of 25 mL/min. For the H₂-TPD experiment, the pre-reduced catalyst surface was kept under H₂ flow at 30 °C for 30 min, followed by purging with Ar for 30 min to remove any physisorbed H₂. Afterward, the desorption was monitored by heating the reactor from 30° to 800 °C at an Ar flow rate of 25 mL/min.

2.3. Thermocatalytic CO2 hydrogenation

2.3.1. CO_2 hydrogenation with H_2

The catalytic tests for CO_2 hydrogenation with H_2 were conducted in a bed stainless-steel flow reactor. Before the reaction, the catalyst precursor was reduced under H_2 flow (30 mL/min) at 300 °C for 1 h. Then, the reactor was fed with a gas mixture of H_2 and CO_2 with a volume ratio of $H_2/CO_2 = 1/1$ (50 mL/min). The reaction products and no converted reactants were evaluated online with a gas chromatograph Agilent Technologies-7890A, equipped with two detectors (one TCD and one FID), one molecular sieve (HP-MOLESIEVE), and 2 columns (one HP-PONA, and one HP-Plot/Q).

2.3.2. CO₂ hydrogenation with water steam

The catalytic tests for CO2 hydrogenation with water steam were carried out in a tubular fixed bed reactor (Ø 5 mm). A scheme of the reaction apparatus is shown in Fig. S1. Before the reaction was carried out, the catalyst precursor was reduced under H2 flow (30 mL/min) at 300 °C for 1 h. In a typical test, CO2 was bubbled into a saturator containing deionized water at 80 °C. CO₂ and saturated water were dragged to the reactor, where the catalyst was located. All piping lines were heated with electrical heating tapes (up to 110 $^{\circ}$ C) to prevent water vapor condensation. For the tests with the mixture of $CO_2 + H_2 + H_2O$, H₂ flow was added to the reactor flow. The flow composition was CO₂/ $H_2/H_2O=1/1/1$ (total flow of 75 mL/min) or $CO_2/H_2O=1/1$ (total flow of 50 mL/min). The temperature of the saturator was determined using Raoult's law and Antoine equation. With Raoult's law, vapor pressure of a desired composition can be find: $y = P_{vap}/P_{total}$, where y is the water steam composition in the gas flow. With Pvap, the temperature of the saturator can be determined via Antoine equation: Pvap = A - B/(C-T), where T is temperature, A = 5.08354, B = 1663.125, and C = 1663.125-45,622 are constants obtained from Bridgeman and Aldrich [44]. The flow of products and unconverted water and CO2 were dragged to the reactor outlet, where the liquid phase was condensed and retained, while the gaseous phase was taken for in-line analysis using chromatography in the gas phase (GC). Ethanol was detected in the condensed liquid phase. The apparatus used to analyze the gaseous and liquid products was a GC Shimadzu 2014, equipped with three detectors (two TCDs and one FID) and 6 columns (two Porapak-Q, one Porapak-N, one MS-13X, one MS-5A, and one Stabilwax). Products selectivity was calculated as following:

$$Selectivity_i(\%) = product_i / \sum product;$$

Productivity_i($\mu mol \cdot g^{-1} \cdot h^{-1}$) = $mols_i/g_{cat} \cdot t$,

where i is the specific product, $g_{cat.}$ is the grams of catalyst used, and t is the reaction time.

The $^1\text{H-NRM}$ analysis was carried out in a AscendTM 600 Bruker spectrometer using 600 MHz of frequency. Typically, 540 μ L of the sample were mixed with 60 μ L of D₂O solution containing 5 mM dimethyl sulfoxide (DMSO) as an internal standard. The spectra were collected with sixteen seconds of relaxation time between the pulses to allow for complete proton relaxation. Water suppression mode was used. MestreNova software was used for data processing.

2.4. Electrochemical methods

2.4.1. Electrodes preparation

The Cu disk electrode (99.99 %, trace metals basis, Mateck) with 1 cm of diameter was polished with alumina suspension followed by electropolishing at $+\ 3\ V$ vs. Cu for $10\ s$ in $66\ \%$ of H_3PO_4 . Ultrapure water was used to rinse the electrode after each procedure. Cu nanocubes were synthesized as described by Roberts, Kulh, and Nilsson [44]. Briefly, the Cu disk, as prepared before, was cycled 4 times at $5\ mV/s$ from $-\ 1.2\ V$ to $+\ 0.9\ V$ vs. RHE in an electrolyte of $0.1\ M$ KHCO3 containing 4 mM KCl. The electrolyte was saturated with CO_2 for at least $15\ min$ before the procedure started.

2.4.2. CO₂ electrochemical reduction test

 ${\rm CO}_2$ electrochemical reduction was performed as described in our previous paper [45]. Typically, ${\rm CO}_2$ was continuously fed at a flow rate of 10 mL/min in a homemade H-cell (made of polyether ether ketone – PEEK) containing 0.1 M KHCO $_3$ as an electrolyte. An anionic exchange membrane (AHO, AGC Inc.) was used to separate the working-electrode compartment from that of the counter-electrode. Each compartment was filled with 10 mL of electrolyte. DSA was used as the counter electrode. All potentials measured against the Ag/AgCl/KCl $_{\rm Sat}$, used as reference electrode, were converted to RHE values according to the Nernst

Equation ($E_{RHE}=E_{Ag/AgCl(KClsat.)}+0.197+0.059$ pH). Biologic SP-300 was the potentiostat used to control the potential of the working-electrode. Liquid and gaseous products were analyzed by HPLC and GC, respectively.

2.4.3. Electrochemically active surface area (ECSA) measurements

ECSA was measured as described by Kanan and co-workers [46,47]. Briefly, the double-layer capacitance was obtained by performing cyclic voltammetry at scan rates of 100, 80, 60, 40, 20, and 10 mV/s in the double layer region (especially between -0.05 V and -0.35 V vs. RHE). 0.1 M HClO₄, Pt mash, and RHE were used as the electrolyte, counter electrode, and reference electrode, respectively. Considering that the number of electrochemically active sites is proportional to the double-layer capacitance value, the roughness factors (RF) of the electrodes were obtained by normalizing the measured capacitances for the flat polycrystalline Cu electrode (conventionally assumed as RF = 1).

3. Results and discussion

3.1. Catalytic performance of ethanol synthesis in the absence or presence of water

Fig. S2 shows the products distribution of the CO_2 hydrogenation ($CO_2/H_2=1$, v/v) on monometallic Cu catalyst at 1 bar (Fig. S2a) and 30 bar (Fig. S2b). The only products detected in the temperature range (160–260 °C) were CO and methanol. Increasing the pressure to 30 bar (Fig. S2b), both methanol and CO production were higher than those at 1 bar (Fig. S2b), but CO was similarly found as the main product. Ethanol and other C_{2+} products were not formed in detectable amounts at both pressures. These results agree with several works that show that C-C coupling is not favored on Cu active sites in CO_2 hydrogenation with CC_2 hydrogenation with

Differently from CO2 hydrogenation in gas-phase, ethanol and other C2+ compounds synthesis from CO2 electroreduction are widely reported in the literature using an aqueous electrolyte saturated with CO2 and applying monometallic Cu as electrode [32,33]. To compare with the results in the gas phase, Cu powder (the same catalyst used in Fig. S2) deposited on a carbon paper was used as the electrode for ethanol synthesis from CO_2 electroreduction at -0.9, -1.0, and - 1.1 V vs. reversible hydrogen electrode (RHE) in a CO₂-saturated KHCO3 (0.1 M) electrolyte. In addition, conventional Cudisk polycrystalline and Cu_{cubes} , synthesized by oxidation-reduction treatment (SEM image is presented in Fig. S3), were also applied to verify the impact of the catalytic surface area and the crystalline phase in the ethanol productivity. CO, ethanol, and ethylene were the main compounds detected for all electrodes. However, here we will present only the results of the liquid ethanol product, which is the focus of this work. Fig. 1 shows that ethanol was formed on all investigated catalysts and potentials applied. Cucubes was the electrode that presented the best catalytic performance for ethanol synthesis due to the preferential exposure of {100} facets in cubes [50]. Notably, {100} facets are essential features for C-C coupling from CO2 in the presence of water [32,33,51–53], and thus it is reasonable that ethanol synthesis was more favored on Cucubes than on polycrystalline Cudisk and Cupowder catalysts. On the other hand, when comparing the results between Cupowder and Cudisk, the best performance in terms of ethanol productivity can be related to the active surface area (inferred here through the roughness factor, RF, shown in Fig. S4 and Table S1), among other factors out of the scope of the present work. The ratio between the RF of Cu_{powder} and that of Cudisk was about 16. Thus, it was expected that a higher amount of ethanol could be synthesized applying $\text{Cu}_{\text{powder}}$ than Cu_{disk} .

The differences observed in the results obtained in electrocatalysis and thermocatalysis are intriguing, suggesting that reaction conditions can lead to different product distributions (e.g., using an aqueous solution in the electrocatalytic system). Considering that ethanol can be synthesized on Cu materials in the CO₂ electrorreduction approach using

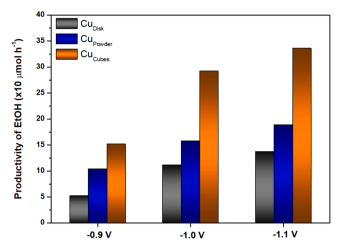


Fig. 1. Productivity of ethanol for Cu_{Disk} (black bars), Cu_{Powder} (blue bars), and Cu_{Cubes} (orange bars) at -0.9 V, -1.0 V, and -1.1 V vs RHE. Reaction conditions: 10 mL of electrolyte (CO_2 -saturated 0.1 M KHCO₃), 1 h of electrolytes.

an aqueous electrolyte (Fig. 1), CO₂ hydrogenation (in gas-phase) was performed as same as performed in Fig. S2a but adding water steam in the feed gas, and the results are shown in Fig. 2. Unlike Fig. S2a, Fig. 2 shows that ethanol could be detected when CO2 hydrogenation with water steam was performed at 1 bar over metallic Cu catalyst. A typical chromatogram obtained from an injection of the condensed liquid (nonconverted water containing liquid products) collected after the flow reactor is shown in Fig. S5a. It is interesting to note that the selectivity to ethanol could reach about 90 % at 170 °C and 180 °C, decreasing drastically at temperatures up to 220 °C. About the productivity, methanol and ethanol augmented when the temperature was increased from 170 $^{\circ}\text{C}$ to 200 $^{\circ}\text{C}$ and decreased at temperatures above 210 $^{\circ}\text{C}.$ It is important to mention that unlike CO2 hydrogenation with H2 at 30 bar where the CO2 conversion can reach about 7% (Fig. S2b), CO2 conversion is thermodynamically limited when the reaction is carried out at 1 bar. Therefore, about 0.1 % is converted for CO₂ hydrogenation with H₂ at 1 bar (Fig. S2a) and less than 0.1% is observed when water is added to the feed stream (Fig. 2). Further discussions about the thermodynamics will be presented in Section 3.3. Different GHSV values were also evaluated -6000, 9000 or 12,000 mL.g⁻¹.h⁻¹ - and the product selectivity and productivity did not change, confirming that the tests were under the kinetic regime.

In parallel with the decline in productivity for alcohols, there was an increase in the productivity for CO. This species is considered one of the

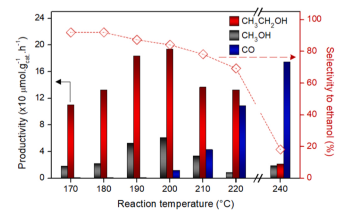


Fig. 2. Productivity of methanol (black bars), ethanol (red bars), CO (blue bars), and selectivity to ethanol between 170 °C and 240 °C. Reaction conditions: $m_{cat.}=500~mg,~CO_2/H_2/H_2O_{(g)}=1/1/1~(75~mL.min^{-1}),~GHSV=9000~mL.g^{-1}.h^{-1},~P=1~bar.$

key intermediaries in proposed reaction mechanisms for ethanol synthesis [32,33,54,55]. Therefore, the presence of adsorbed CO is essential for the synthesis of ethanol. However, the formation of CO as a reaction product means that CO produced in situ by the hydrogenation of CO_2 is desorbing rather than remaining adsorbed on the surface and being further hydrogenated for the formation of ethanol. In this work, $180-200\,^{\circ}C$ is the range of temperature that most favors the formation of ethanol. It is hard to compare the catalytic activity showed in this work with previous results in the literature as the reaction conditions applied here are not commonly used: 1 bar and adding water in the feed stream. However, a summary of reaction conditions, ethanol selectivity, CO_2 conversion, etc., of several recent works focused on ethanol formation from CO_2 hydrogenation are presented in Table S2.

The role of water as a hydrogen donor was clarified by carrying CO_2 hydrogenation with water steam in the absence of H_2 flow, and results are shown in Fig. S6. Interestingly, the same trend of methanol, ethanol, and CO productivity was observed. To the best of our knowledge, here we show for the first time that ethanol can be synthesized at atmospheric pressure, applying metallic Cu catalyst and using only CO_2 and water as reactants in a thermocatalytic process. A thermodynamic evaluation of the reaction and the catalyst deactivation will be discussed later in Section 3.3. Ethanol productivity in the absence of H_2 was slightly lower than that of the reaction with H_2 , which indicates that the ethanol synthesis is mainly produced via $CO_2 + H_2O$. More insights about the water involvement in the synthesis of ethanol will be shown in Section 3.2.

Inspired by the electrocatalytic results (Fig. 1), CO₂ hydrogenation with water steam (without H2 flow) was also performed in Cu catalysts with different surface areas and Cucubes. This study makes it possible to understand the influence of the surface area and the crystalline phase in ethanol synthesis in a thermocatalytic process as observed in electrocatalysis. Fig. 3 shows the ethanol productivity at 170 °C, 190 °C, and 210 °C by applying Cu catalysts obtained from (i) direct calcination of Cu(NO₃)₂, named Cu_{calc.}; (ii) precipitation of Cu(OH)₂ followed by calcination, named Cuprec.; and (iii) controlled synthesis of Cu2O to produce Cu nanocubes, as described by Chang et al. [43]. Before the reaction was carried out, Cu_xO was reduced in situ to Cu at 300 °C with H₂ for 1 h. Ethanol productivity decreased in the following order: $\text{Cu}_{\text{cubes}} > \text{Cu}_{\text{prec.}} > \text{Cu}_{\text{calc}}.$ The differences can be partially justified by the material's surface areas. CuxO surface area was found to be 27 m²/ $g_{cat.}$ for Cu_{cubes} , 20 m²/ $g_{cat.}$ for $Cu_{prec.}$, and 11 m²/ $g_{cat.}$ for $Cu_{calc.}$ Higher surface area may result in a higher amount of exposed Cu sites for CO₂ hydrogenation, which can lead to higher ethanol production. When the ethanol productivity was normalized by Cu_xO surface area (m²Cu) rather than of catalyst mass (g_{cat.}) (Fig. 3b), it became clear that Cu_{prec.} and Cucalc, show similar results in terms of ethanol productivity. These data show that, for Cuprec and Cucalc, ethanol productivity depends directly on the CuxO surface area. On the other hand, ethanol productivity on Cucubes remains higher than on Cuprec, and Cucalc, even considering the surface area of the catalysts, which suggests that an additional feature in the Cucubes is acting to improve its catalytic performance. SEM images of Cucubes synthesized in this work are presented in Fig. S7. It is possible to see regular nanocubes with 47 \pm 5 nm of edges, like those produced by Chang et al. [43], who also showed by HR-TEM that these nanocubes are {100} facets. Further surface characterization of the Cu nanocubes can be found in the work of Chang et al. [43]. Therefore, as evidenced in the CO₂ electroreduction approach, $\{100\}$ facets preferably exposed in the Cu_{cubes} are also crucial in the hydrogenation of CO2 with water steam. Similar to what was observed in Fig. 2 in electrocatalysis, ethanol synthesis from CO₂ and water steam is both area- and crystalline structure-sensitive.

3.2. Water involvement in the ethanol synthesis

Since the ethanol productivity is relatively low using only ${\rm CO_2}$ and water in the thermocatalytic process, a natural question concerns the

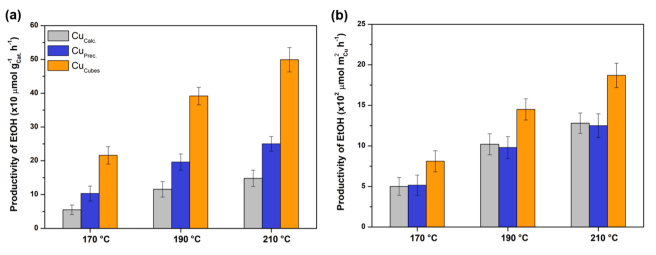


Fig. 3. Productivity of ethanol (a) per $g_{cat.}$ and (b) per m^2_{Cu} for $Cu_{Calc.}$ (gray bars), $Cu_{Prec.}$ (blue bars), and Cu_{Cubes} (orange bars) at 170 °C, 190 °C, and 210 °C. Reaction conditions: $m_{cat.} = 500$ mg, $CO_2/H_2O_{(g)} = 1/1$ (50 mL.min⁻¹), GHSV = 6000 mL.g⁻¹.h⁻¹, P = 1 bar.

possibility of the ethanol synthesis occurring by the reaction between the CO_2 and hydrogen chemisorbed on Cu , from the previous reduction step of $\mathrm{Cu}_x\mathrm{O}$ to Cu with H_2 . To investigate this possibility, H_2 -TPD was performed on the Cu catalyst. In this analysis, no H_2 desorption was detected (Fig. S8a) after the reduction step, indicating that no residual hydrogen was chemically adsorbed on the Cu surface when the CO_2 hydrogenation reaction started. To confirm the reliability of these results, Cu^+ from pre-step oxidation of Cu with $\mathrm{N}_2\mathrm{O}$ was studied by H_2 -TPR. Fig. S8b shows hydrogen consumed in the Cu^+ to Cu^0 reduction process, attesting to the detection capability of the employed equipment at the investigated scale.

The involvement of water in the CO₂ hydrogenation was studied by 1 H NMR. CO $_{2}$ hydrogenation was performed with H $_{2}$ O and, additionally, with a mixture of D₂O/H₂O steam at 190 °C, and the condensed liquid products were analyzed by ¹H NMR. It is essential to point out that before the reaction was carried out and to prevent misinterpretation of data, the natural exchange between hydrogen and deuterium in the ethanol molecule was investigated by preparing a standard ¹H NMR sample with 1 mmol/L in D2O, which was aged for three days. The results show that triplet and quadruplet generated by CH3 and CH2 coupling, respectively, were not affected by D2O, but the singlet generated by OH group (expected at ~2.61 ppm) presents an exchange to form OD in the presence of D₂O (Fig. S9 and Supplementary Note 1). So, natural exchange in CH3 and CH2 groups in solution between deuterium and the hydrogen from ethanol formed from CO2 hydrogenation will not be considered here. Fig. 4 illustrates the multiplets and singlets generated by the couplings of the three non-equivalent hydrogen atoms present in ethanol and compares the spectra of condensed products of H₂O (black lines) and D₂O/H₂O containing reaction (blue lines). The characteristic triplet, generated by the coupling of hydrogen in CH3 with the two hydrogens in CH2, decreased its intensity when the reaction was carried out in the presence of D₂O. A doublet emerged, indicating the partial substitution of hydrogen by deuterium in the CH2 group. The same effect was observed in the quadruplet generated by hydrogen in CH2 with the three hydrogens in CH₃, indicating the partial substitution of hydrogen by deuterium in the CH₃ group. As mentioned before, the complete disappearance of the singlet, generated by the hydroxyl group of ethanol, can not be consistently attributed to substitution during reaction due to the natural exchange observed. The signal decreasing and the appearance of lower multiplicity peaks in CO2/H2O/D2O reaction in relation to CO2/H2O indicate the presence of partially deuterated molecules in two nonequivalent hydrogen positions of ethanol (CH3 and CH2), attesting that water is responsible for providing atomic hydrogen during the reaction. Additionally, as shown before, when the reaction was carried out

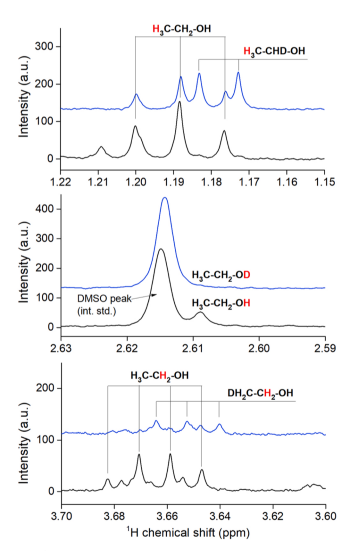


Fig. 4. 1 H NMR spectra of condensed products from CO₂ hydrogenation in the presence of H₂O (black lines) and H₂O/D₂O (blue lines). Reaction conditions: T = 190 $^{\circ}$ C, m_{cat.} = 500 mg, CO₂/H₂O_(g) = 1/1 (50 mL/min), P = 1 bar.

in a flow of $CO_2/H_2O/H_2$ (Fig. 2), no considerable increase in the production of ethanol was observed in relation to the use of CO_2/H_2O flow (Fig. S6), revealing that water is the primary hydrogen source to produce ethanol at the investigated conditions.

3.3. Thermodynamics discussion

The thermodynamic is one of the key questions about ethanol synthesis from CO₂ and water. Considering the global reaction: 2CO_{2(g)} + 3 $H_2O_{(g)}\to C_2H_5OH_{(g)}+3$ $O_{2(g)},$ it results in a reaction with high endothermic energy ($\Delta H=1277.3$ kJ/mol). Under the reaction conditions tions applied in this work, ethanol might not be formed from the direct hydrogenation of CO2 with water steam. In fact, some experimental insights indicate that ethanol can be formed from a different path. After the reduction step, it was possible to see by bare eyes that the catalyst was fully brownish, characteristic of metallic copper, while the catalyst was majority blackish after the reaction with water, indicating surface oxidation during the reaction. The material surface oxidation was also seen experimentally by the deactivation of the catalyst over time, as shown in Fig. 5a. Specifically, using a fresh catalyst sample, the ethanol productivity was practically suppressed after 3 h. However, when the same sample was reactivated, by reducing it again with H₂ at 300 °C for 1 h, ethanol synthesis was again detected (schematically exemplified in Fig. 5b). However, a lower productivity was observed than the fresh catalyst was used. This decay in productivity comparing both situations can be related to reducing the surface area. The surface area of the metallic Cu was measured to be 8.7 m²/g, while the same material reactivated was $2.2 \text{ m}^2/\text{g}$, about 4 times smaller than the fresh sample. The same behavior was also observed for the second reactivation, although the decay in the ethanol productivity was smaller. This can again be explained by the surface area. The metallic surface area was measured to be 1.8 m²/g, a value close to the measured before, so a strong decay is not expected as seen for the first reactivation. For the third reactivation, similar ethanol productivity was observed compared to the second reactivation, which was expected as the same metallic surface area was measured.

The deactivation observed in Fig. 5a indicates that Cu^0 is oxidized over time to Cu^{2+} and/or Cu^+ by water reduction, $\text{Cu} + \text{H}_2\text{O} \leftrightarrow \text{Cu}\text{O} + \text{H}_2$, $2\text{Cu} + \text{H}_2\text{O} \leftrightarrow \text{Cu}_2\text{O} + \text{H}_2$. The oxidation of Cu by the presence of water leading to its deactivation has been studied by different groups. For example, Prašnikar and Likosar showed that the water formed in-situ from CO_2 hydrogenation to methanol increased the Cu deactivation rate [56,57]. In the same direction, Chen *et. al* showed by in-situ DRIFT and DFT calculation that the reduction of ethyl acetate on Cu is strongly affected by water adsorption [58]. More specifically, the active species

of Cu^+ on the surface has preferred the adsorption of $\mathrm{H}_2\mathrm{O}$ over the ethyl acetate molecule leading to a rapidly decreasing in the conversion ratio of ethyl acetate. At this moment, we do not have yet evidence whether Cu^0 , copper oxides or a mixture of Cu^0 /copper oxides is the active site for the ethanol formation in the catalytic reaction that we presented in this work. It is well known that Cu^+ is an important active site for methanol synthesis from CO_2 hydrogenation [59]. So, Cu^+ probably also plays an important role in the formation of ethanol. In this way, $\mathrm{H}_2\mathrm{O}$ might also adsorb strongly on the catalyst surface, as observed by *Chen* et al. [58], favoring its deactivation.

With the oxidation of Cu metallic, hydrogen is formed in-situ by the water reduction which might drastically decrease the energy needed to form ethanol, turning the reaction exothermic. Specifically, the ethanol synthesis considering H_2 as the reducing agent $(2\text{CO}_{2(g)}+6\ H_{2(g)}\to \text{C}_2\text{H}_5\text{OH}_{(g)}+3\ \text{H}_2\text{O}_{(g)})$ has a $\Delta H=-173.5\ \text{kJ/mol}.$ So, combining the total oxidation reaction of Cu and the ethanol synthesis from CO $_2$ and H_2 , we can write the global reaction for ethanol synthesis from CO $_2$ and water steam applying metallic copper as the catalyst, as shown in Eq. 3.

$$6Cu_{(s)} + 6 H_2O_{(g)} \rightarrow 6CuO_{(s)} + 6 H_{2(g)} \Delta H = 507 \text{ kJ/mol}$$
 (1)

$$2CO_{2(g)} + 6 H_{2(g)} \rightarrow C_2H_5OH_{(g)} + 3 H_2O_{(g)} \Delta H = -173.5 \text{ kJ/mol}$$
 (2)

$$6Cu_{(s)} + 2CO_{2(g)} + 3 H_2O_{(g)} \rightarrow C_2H_5OH_{(g)} + 6CuO_{(s)} \Delta H = 333 \cdot 5 \text{ kJ/mol(3)}$$

Some important assessments can be done by looking at the global reaction. Firstly, the ΔH is still high compared to other reactions such as the traditional methanol synthesis from CO2 hydrogenation ($\Delta H = -49.5 \ kJ/mol$), which explains why the ethanol productivity is rather low at the reaction conditions applied in this work. Secondly, as shown in Fig. 5b, a stepwise process is proposed since Cu acts as both catalyst and reactant to produce ethanol and in-situ H_2 , respectively. The active sites in the metallic copper surface are being saturated due to adsorption of oxygen from water, forming a shell of CuxO, which is responsible for deactivating the catalyst very quickly. The oxygen can be removed by successive steps of H_2 feeding between reaction steps.

In an electrochemical system, the deactivation by the oxidation of the Cu electrode is not so evident because the potentials normally applied for CO $_2$ reduction to ethanol (<-0.7 V) are considerably more negative than those for Cu $^+$ and Cu $^{2+}$ reduction (Cu $^+$ + e \leftrightarrow Cu $_{(s)}$, E =+0.52 V; Cu $^{2+}$ + 2e \leftrightarrow Cu $_{(s)}$, E =+0.34 V). To optimize the stability of the Cu in the thermocatalytic CO $_2$ hydrogenation with water steam, increasing the H_2/H_2O ratio could be an interesting strategy to enhance the reducibility of Cu $_x$ O. However, H_2 -TPR shows that Cu oxide powder reduces only at temperatures over 250 °C (Fig. S10) and thus H_2 will not be able to reduce Cu $_x$ O in the temperature range for ethanol synthesis

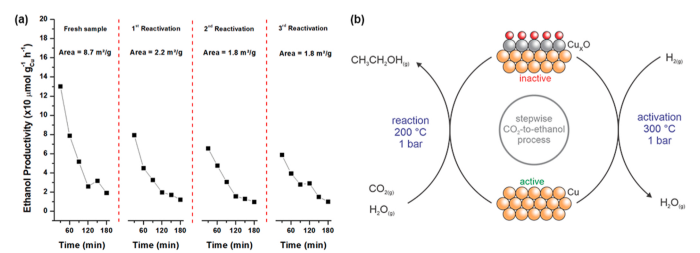


Fig. 5. (a) Ethanol productivity over time. Dashed red lines mean that a new reactivation of the Cu catalyst with H₂ at 300 °C was performed. (b) Proposed stepwise process model for CO₂-to-ethanol conversion over Cu catalyst.

(between 170 and 220 °C). Over 220 °C, better Cu reduction could be achieved but ethanol production would be suppressed, as shown in Fig. 2. Cu supported on oxides which lead for lower Cu temperature reduction must be considered for future studies. Therefore, water is a key component that acts both positively and negatively for ethanol synthesis from the hydrogenation of CO_2 with water steam. Finally, a natural question to be understood is why ethanol can be produced by adding water steam in the feed gas, but it is not formed when only CO_2 and H_2 are applied (Fig. S2). CO-DRIFTS experiments described in the next section give us an insight into the importance of the water steam in the reaction system.

3.4. CO-Drifts

Results have shown that ethanol can be synthesized from CO_2 hydrogenation in the presence of water steam, although the fundamental role of water in the process is unclear. CO-DRIFTS tests were carried out to understand how the adsorption of CO is influenced by the presence and absence of water steam. CO was chosen as a probe molecule since it is considered a key intermediate in the CO_2 hydrogenation and the reaction mechanism for C_{2+} synthesis. Previous tests (results not shown here) at standard conditions (15 °C) with monometallic Cu and catalysts with a high Cu loading (Cu/SiO₂, Cu/Al₂O₃, and Cu/CeO₂ with molar ratio of Cu/(SiO₂ or Al₂O₃) = 1/1) were carried out, and no signals referring to the CO bands were detected after the purging step with N_2 . Therefore, monometallic Cu or catalysts with a high Cu loading could not be employed at the investigated conditions, requiring tests at very low temperatures [60]. However, at temperatures below 0 °C, water steam would condense, and these tests could not be performed.

On the other hand, previous works show that the application of a catalyst with low Cu loading supported on an oxide, such as the $2.5\%\text{Cu}/\text{CeO}_2$ compound, allows well-dispersed Cu⁺ nanoparticles to be formed during the catalyst synthesis and the chemisorption of CO on Cu⁺ (Cu⁺-CO) remains stable even after purging with an inert gas (N₂, Ar or He) [60–62]. As discussed before, Cu⁺ is pointed out as active site for the methanol synthesis from CO₂ hydrogenation and it is likely that Cu⁺ is also partially formed during the Cu oxidation via water reduction in the CO₂ hydrogenation with water steam. Therefore, for the spectroscopic tests, Cu/CeO₂ (2.5 molar% of Cu in CeO₂) was chosen as a model catalyst in order to increase the sensibility of CO adsorption on the Cu active phase in the presence and absence of water steam.

Fig. 6 shows the CO-DRIFTS results. Before CO adsorption, the CuO/CeO $_2$ catalyst was reduced in situ to Cu/CeO $_2$ with a flow of 30 mL/min

of H2 at 300 °C for 1 h. The background was obtained at 15 °C after 15 min of purging with N2 after the reduction process. The spectrum shown in Fig. 6 consists of the profile after the background subtraction and the bands are related to the CO adsorption. Linear CO was detected at 2119 cm⁻¹ with shoulders at 2138 cm⁻¹ and 2095 cm⁻¹, in accord with the literature [60,63]. The bands were deconvoluted with Voigt adjustment, and the result is shown in Fig. 6b. The total fit (dashed curve in red) was adequately adjusted to the experimental signal (curve in black), with the residual signal (curve in green) representing the difference between the experimental curve and the fit being null. This adjustment made it possible to identify three well-defined bands of adsorbed CO, centered at 2139, 2120, and 2105 cm⁻¹. The main band at 2120 cm⁻¹ is attributed to Cu⁺-CO species, as shown by many authors [60,61,64,65]. In addition, Cu^0 –CO species can also appear at the same wavenumber as Cu⁺-CO and these two species can be distinguished by their stability, being CO adsorbed on Cu⁰ easily removed during the purging [60,61]. In the present study, since the 2120 cm⁻¹ band was highly stable to N₂ purge, it is more likely to be related to Cu⁺-CO. The shoulders at 2139 cm⁻¹ and 2105 cm⁻¹ are pointed out to be dicarbonyl species of Cu⁺-(CO)₂, characterized by symmetrical and antisymmetrical vibration modes [61,66].

To investigate the adsorption of CO on Cu/CeO₂ in the presence of H₂ or H₂O steam, the same procedure performed previously was applied, but with CO adsorbing together with either H2 or H2O steam and the spectra after 15 min of purging with N2 are shown in Fig. 7a. The spectrum obtained by exposing the catalyst only to the CO flow was also added to the figure as a reference. No significant difference is identified by comparing the profiles of the Cu⁺-CO and Cu⁺-(CO)₂ bands for the CO adsorption on Cu/CeO2 with and without hydrogen in the medium (red line and black line, respectively). On the other hand, in the presence of water (blue line), it can be observed that the adsorption bands referring to Cu⁺-CO and Cu⁺-(CO)₂ shifted to smaller wavenumbers. Fig. 7b, c, and d show the bands evidenced in the experiments with CO, CO + H₂, and CO + H₂O, respectively, deconvoluted by Voigt adjustment. It becomes clear that the Cu⁺-CO and Cu⁺-(CO)₂ bands, located at 2120, 2139, and 2105 cm⁻¹, respectively, in the tests with CO and CO + H₂, moved to 2110, 2120, and 2098 cm⁻¹, respectively, in the test with CO + H₂O. This shift means that lower energies were needed to promote the vibrations of the C-O stretching of the adsorbate on the catalyst surface due to a weakening of the carbon-oxygen bond referring to the CO adsorbed on Cu. Simultaneously with the weakening of the carbon-oxygen bond, the carbon-metal bond is strengthened. In this way, water seems to stabilize CO on the Cu surface. These results may

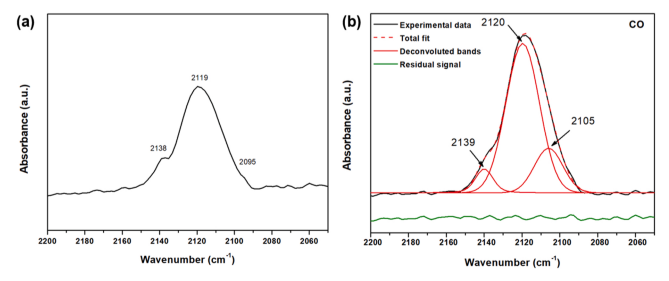


Fig. 6. (a) DRIFTS spectrum of CO adsorption on Cu/CeO₂ at 15 °C and atmospheric pressure followed by N₂ purge. (b) Deconvolution of the CO band by Voigt adjustment.

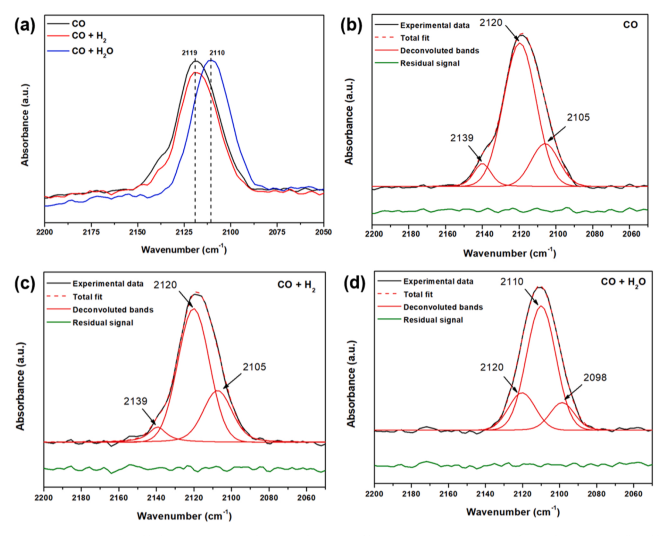


Fig. 7. (a) DRIFTS spectrum of CO (black curve), $CO + H_2$ (red curve), and $CO + H_2O$ (blue curve) on Cu/CeO_2 at 15 °C and atmospheric pressure followed by N_2 purge. Deconvolution of the CO_{ad} band by Voigt adjustment when the tests were carried out with (b) CO; (c) $CO + H_2$; and (d) $CO + H_2O$.

explain why C_2 compounds are favored in the hydrogenation of CO_2 with water steam, whereas only C_1 compounds are observed when H_2 is used instead of water. As CO is considered a key intermediate in the

synthesis of ethanol from CO₂, the strengthening of the carbon-metal bond, observed only in the CO adsorption tests with water steam, lead to maintaining CO adsorbed on the surface and, therefore, it is more

i) Water adsorption

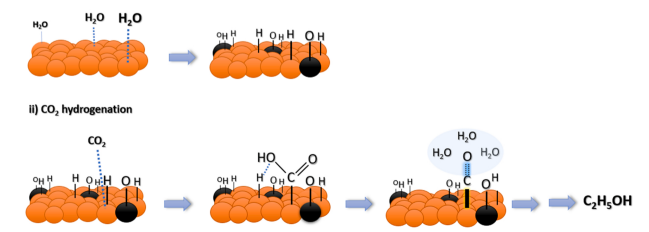


Fig. 8. Proposed scheme of (i) Water adsorption and (ii) CO₂ hydrogenation with water steam on Cu catalysts. The CO₂ adsorption is represented here on Cu and/or Cu⁺.

likely that it continues to react rather than desorbing as a product.

A scheme representing the first steps of the reaction mechanism for ethanol synthesis from CO₂ and water steam is presented in Fig. 8, taking into account the catalytic results, thermodynamic discussion, and the DRIFTS tests. It is important to mention that CO2 can adsorb in different ways depending on the catalyst and reaction conditions [67-69]. *HCOO is considered the main intermediate for methanol synthesis (called formate pathway) and it is believed that CO is not formed from this pathway [70]. *COOH is reported to be a common intermediate for CO formation as in the presence of H^* , $\mathrm{H}_2\mathrm{O}$ and CO are formed from the rupture of CO-OH [27,70]. The dissociative CO₂ adsorption (*CO+*O) is also reported to be a route for CO formation but *O must be consumed by *H to form *OH or H₂O and/or react with another *O to form O₂ [71]. Both dissociative CO2 adsorption and *COOH may (simultaneously) happen, but as in our tests the system has a high concentration of water, and water can easily adsorb and oxidize the Cu surface (shown in Fig. 5), high concentration of H* will also be present on the Cu surface. For a representative scheme, the *COOH pathway was chosen here but further investigation must be done. In summary, one step consists of water adsorption and splitting on the Cu surface followed by its oxidation. Simultaneously, CO₂ is adsorbed on Cu and/or Cu⁺ and further hydrogenated to CO. As shown by DRIFTS tests, a strengthening of the carbon-metal interaction is favored in the presence of water (represented by a thicker Cu-C bond). Finally, the most strongly adsorbed CO is further hydrogenated to ethanol.

4. Conclusions

Cu catalysts are not active for ethanol synthesis from CO₂ hydrogenation under standard conditions ($CO_2 + H_2$), although the opposite is observed in an electrochemical system (CO₂ + H₂O). Inspired by the electrochemical results, we showed that ethanol could be synthesized thermocatalytically by adding water steam in the feed gas (CO2 + H2 + H₂O) with selectivity of 84 % and productivity of \sim 2 μ mol.g_{cat}⁻¹.h⁻¹. When only CO₂ and water steam were used, the same trend for ethanol synthesis was observed. This reaction showed to be both area- and crystalline structure- sensitive. Higher surface area and {100} facets showed better catalytic performance for ethanol synthesis. Applying deuterated water steam, it was possible to identify by ¹H NMR that water is acting as a hydrogen donor for the synthesis of ethanol. The direct synthesis of ethanol from CO₂ shows a high enthalpy value. Hence, the most likely reaction consists in the in-situ hydrogen formation by the oxidation of metallic copper while the water reduction occurs, followed by the hydrogenation of CO_2 into ethanol: 1) $6Cu_{(s)} + 6H_2O_{(g)} \rightarrow$ $6CuO_{(s)} + 6 H_{2(g)}$; 2) $2CO_{2(g)} + 6 H_{2(g)} \rightarrow C_2H_5OH_{(g)} + 3 H_2O_{(g)}$. The low ethanol productivity evidenced in this work is explained by the high enthalpy ($\Delta H = 333.5 \text{ kJ/mol}$) of the global reaction: $6Cu_{(s)} + 2CO_{2(g)}$ $+ 3 H_2O_{(g)} \rightarrow C_2H_5OH_{(g)} + 6CuO_{(s)}$. By DRIFTS analysis, a strengthening of the carbon-metal interaction was observed only for the CO adsorption in presence of water steam, which may be one of the reasons why ethanol is favored in the CO₂ hydrogenation with water steam.

The approach used in this work for ethanol synthesis from CO_2 and water steam is far from a real application since the ethanol productivity and catalyst stability are rather low. However, this work opens new possibilities for future research related to CO_2 hydrogenation, considering the opportunity for high selectivity to higher alcohols by water enhancement. Using a feed gas mixture of CO_2 , H_2 , and H_2O over a catalyst more resistant to the water reduction (e.g., copper supported in oxides and multi-metallic materials) may be an interesting approach since better catalyst stability and the water-enhancement in carbon-metal strengthening can be both achieved simultaneously.

CRediT authorship contribution statement

Alisson H. M. da Silva: Conceptualization, Formal analysis, Investigation, Data curation, Writing – original draft, Luiz H. Vieira:

Investigation, Cássia S. Santana: Investigation, Marc T. M. Kopper:review & editing, Supervision, Elisabete M. Assaf:- Resources, review & editing, Supervision, Janaina F. Gomes: Conceptualization, Data Curation, Resources, Writing – review & editing, Supervision, José M. Assaf: Conceptualization, Recourses, Writing – review & editing, Supervision, Funding acquisition.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.apcatb.2022.122221.

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